1



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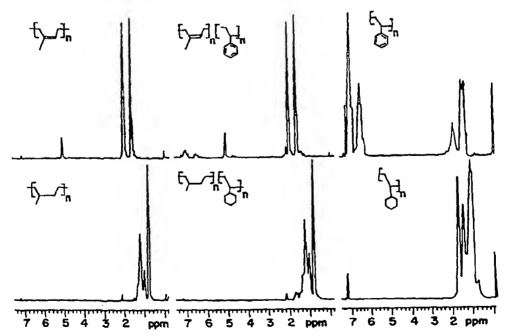
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(54) Title: METHOD FOR HYDROGENATING POLYMERS AND PRODUCTS THEREFROM



(57) Abstract

This invention relates to a method for hydrogenation of poly(alkenyl aromatic) polymers or poly(alkenyl aromatic)/polydiene block copolymers that provides hydrogenated polymers with 99.5 % or greater unsaturation and an MWD of less than about 3. Hydrogenated polymers and adhesives comprising the hydrogenated polymers and tackifier are also provided.

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. , WO 94/21694 PCT/US94/02720

- 2 -

the reactions which may be performed on some of these block copolymers is a hydrogenation reaction wherein the unsaturations within the blocks are removed. For example, in a styrene-isoprene-styrene block copolymer a homogeneous catalyst may be utilized to selectively hydrogenate the unsaturations in the isoprene to produce what is essentially an ethylene propylene block while leaving the styrene blocks saturated. This particular configuration has found many uses and is known commercially as KRATON G.

Hydrogenation of the isoprene and the polystyrene blocks in a multiblock copolymer has also been reported in the art. U.S. Patent No. 3,333,024 to Haefele 15 discloses that a styrene-isoprene-styrene block copolymer having a combined styrene content of 22% by weight can be hydrogenated by suspending the block copolymer in cyclohexanes and contacting it with a catalyst of nickel on kieselguhr at a pressure of 500 20 psig and at a temperature of 145-55°C for thirteen hours. The resulting hydrogenated block copolymer had 98% hydrogenation of the polystyrene blocks. The block copolymers were reported to have improved tensile properties over the unhydrogenated block copolymers. 25 However, in Chapter 14 of Thermoplastic Elastomers, a comprehensive review, Ed. N.R. Legge, et al., Hanser Publishers, New York, 1987, this class of hydrogenated block copolymers was described as being generally poor. These hydrogenated materials are generally described as 30 having an interaction parameter so severely reduced that at only slightly elevated temperatures the polymer loses all strength and appears to be homogeneous mixture. Thus, the good tensile strength of U.S. 3,333,024 appears to be limited to a low temperature 35 end use range.

WO 94/21694

- 3 -

The hydrogenation of polymers is a known technique. There are two typical techniques, homogeneous and heterogeneous that have been shown effective for obtaining saturated polymers. 5 Homogeneous hydrogenation yields high conversions, however, this method sometimes causes chain scission and often leads to metal contamination due to a catalyst extraction difficulties. Heterogeneous hydrogenation yields high conversions and in certain 10 cases is more tractable than homogeneous hydrogenation due to minimal chain scission and no metal contamination and the ease of catalyst extraction. Furthermore, supported heterogenous transition metal catalysts are recyclable with no byproducts unlike 15 homogeneous extraction products which generate large amounts of waste product. Heterogeneous hydrogenation has been used to saturate many types of polydienes including 1,4 and 3,4-poly(isoprene) and 1,2 and 1,4polybutadiene. (Zhongde, X; et al., Macromolecules, 20 1983, 16, 925.)

Thus, hydrogenated block copolymers having good end-use properties such as high order/disorder transition temperatures, (ToDT) or high glass

25 transition temperatures (Tg's) and methods to make them are desirable.

Summary of the Invention

30 This invention relates to a method for hydrogenating polymers comprising contacting the copolymer with a metal catalyst on an Group 1 or 2 metal salt support. This invention further relates to hydrogenated polymers having greater than 99.5% saturation and a molecular weight distribution (Mw/Mn) of less than about 3, preferably less than about 2,

' ' WO 94/21694 PCT/US94/02720

- 4 -

even more preferably less than about 1.5. In another aspect, this invention relates to various compositions containing and varying uses for the hydrogenated polymers such as adhesives.

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Brief Description of the Drawings

Figure 1 is proton NMR spectra showing polystyrene polyisoprene diblock copolymer before and after

10 hydrogenation. (hydrogenated polymer on the bottom)

Description of Preferred Embodiments

This invention relates to a method for

15 hydrogenating unsaturated polymers, preferably
poly(alkenyl aromatic) homo- or copolymers or
poly(alkenyl aromatic)/polydiene block copolymers
comprising contacting, under reaction conditions, the
unsaturated polymer with a metal catalyst on an Group 1

20 or 2 metal salt support and recovering the hydrogenated
polymer having greater than 99.5% saturation and a
molecular weight distribution of less than about 3.

This invention further relates to hydrogenated

25 poly(alkenyl aromatic) homo- and copolymers and
hydrogenated block copolymers comprising hydrogenated
poly(alkenyl aromatic)/hydrogenated polydiene blocks
having greater than about 99.5% unsaturation,
preferably 99.75%, and a Mw/Mn of less than about 3,

30 preferably less than about 2, even more preferably less
than about 1.5.

In particular, this invention provides for hydrogenated poly(alkenyl ar matic), preferably hydrogenated polystyrene (also called poly(vinylcyclohexane), hydrogenated syndiotactic

• . WO 94/21694 PCT/US94/02720

- 5 -

poly(alkenyl aromatic), preferably hydrogenated syndiotactic polystyrene, hydrogenated polystyrene/hydrogenated polydiene block copolymers, preferably hydrogenated (polystyrene-polyisoprene-polystyrene) block copolymers and hydrogenated (polystyrene-polybutadiene-polystyrene) block copolymers having greater than 99.5%, preferably 99.75%, unsaturation and a molecular weight distribution of less than about 3, preferably less than about 2, even more preferably less than about 1.5.

This invention further relates to adhesives comprising tackifiers, hydrogenated polymers of this invention and optional additives known to those of ordinary skill in the art.

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For the purposes of this application and the claims appended thereto the term block copolymers is herein specifically defined to include diblocks, 20 triblocks, multiblock, tapered block and star block copolymers, and the term copolymer is defined to mean polymers of two or more different monomers.

In a preferred embodiment the homo- and copolymers
of the poly(alkenyl aromatic) include random,
isotactic, syndiotactic and amorphous homopolymers and
copolymers of any alkenyl aromatic hydrocarbon,
preferably where the alkyl has 1 - 20 carbon atoms,
even more preferably 2-8 carbon atoms, and the aromatic
is mono or polycyclic and can be substituted or
unsubstituted, preferably with one or more alkyls each
independently containing 0 to 20 carbon atoms, even
more preferably 0 to 8 carbon atoms. Preferred
polymers include polystyrene, syndiotactic polystyrene,
p ly(para alkylstyrene) & syndiotactic poly (para
alkylstyrene) where the alkyl has 1 to 20 carbon atoms,

preferably 1 - 8 carbon atoms, even more preferably 1
carbon atom.

Stereospecific alkenyl aromatic homo- and

copolymers, particularly syndiotactic poly (alkenyl aromatic)s, even more particularly, syndiotactic polystyrenes, can be made by the processes disclosed in US 4924018, US 5093295, US 5070160, JP 62158709-A (pub. 7-14-87), JP 04258645-A (pub. 9-14-92), EPA 423100-A (pub. 4-17-91) which are incorporated by reference herein.

The polydienes in the starting block copolymers can be any polydiene, preferably polymers of a C4 to C20 diene, preferably C4 to C10. Another preferred embodiment contains conjugated polyenes, conjugated dienes or alpha-omega dienes. It is known that when an alpha-omega diene or a conjugated diene is added to a growing polymer that it can be added by different 20 mechanisms of addition. For example 1,4-butadiene can be added by 1,2 addition or by 1,4 addition. Depending on the final use of the hydrogenated block copolymer, varying amounts of differently added monomer will be desirable in the starting block copolymer. For 25 example, in styrene/butadiene block copolymer applications where flexibility is of greater importance a significant amount of 1,2 added butadiene would be desirable since it hydrogenates to something like an ethylene/butene polymer. 1,4 added polybutene hydrogenates to something like ethylene polymer and would be much stiffer, and therefore would be preferable for applications requiring greater stiffness. It is within the scope of the invention that the polydiene block may comprise anywhere from 0 to 100% 1,2 added monomer and from 100% to 0% other forms of added monomer. For polybutene, up to 100% of

· . · WO 94/21694 PCT/US94/02720

- 6 -

preferably 1 - 8 carbon atoms, even more preferably 1 carbon atom.

Stereospecific alkenyl aromatic homo- and

copolymers, particularly syndiotactic poly (alkenyl aromatic)s, even more particularly, syndiotactic polystyrenes, can be made by the processes disclosed in US 4924018, US 5093295, US 5070160, JP 62158709-A (pub. 7-14-87), JP 04258645-A (pub. 9-14-92), EPA 423100-A (pub. 4-17-91) which are incorporated by reference herein.

The polydienes in the starting block copolymers can be any polydiene, preferably polymers of a C4 to 15 C₂₀ diene, preferably C₄ to C₁₀. Another preferred embodiment contains conjugated polyenes, conjugated dienes or alpha-omega dienes. It is known that when an alpha-omega diene or a conjugated diene is added to a growing polymer that it can be added by different 20 mechanisms of addition. For example 1,4-butadiene can be added by 1,2 addition or by 1,4 addition. Depending on the final use of the hydrogenated block copolymer, varying amounts of differently added monomer will be desirable in the starting block copolymer. For 25 example, in styrene/butadiene block copolymer applications where flexibility is of greater importance a significant amount of 1,2 added butadiene would be desirable since it hydrogenates to something like an ethylene/butene polymer. 1,4 added polybutene 30 hydrogenates to something like ethylene polymer and would be much stiffer, and therefore would be preferable for applications requiring greater stiffness. It is within the scope of the invention that the polydiene block may c mprise anywhere from 0 35 to 100% 1,2 added monomer and from 100% to 0% other forms of added monomer. For polybutene, up to 100% of

· WO 94/21694 PCT/US94/02720

- 7 -

the monomer may be added by 1,2 addition, preferably up to 75% The amount of monomer added by a particular form of addition may be influenced by the isomeric configuration of the monomer or by the catalyst used to 5 make the blocks.

Examples of preferred polydienes include, but are not limited to, homopolymeric and copolymeric polymers of butadiene, pentadiene, hexadiene, heptadiene, octadiene, nonadiene, isobutadiene, isopentadiene, isohexadiene, isohexadiene and isoprene.

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The poly(alkenyl aromatics) in the starting block copolymers may be polymers of any alkenyl aromatic hydrocarbons specifically alkyl aromatic hydrocarbons wherein the aromatic may either be monocyclic or polycyclic, substituted or unsubstituted, preferably with one or more alkyls having 0 to 20 carbon atoms, preferably 0 to 8 carbon atoms. and the alkyl has 1 to 20 carbons, preferably 2 to 8 carbons. Preferred species include styrene, methylstyrene, alpha methyl styrene, para methyl styrene, para dimethlstryene, butlystyrene, para t-butyl styrene, vinylxylene, ethylvinylxylene, isopropylstyrene, vinylnapthalene and the like.

The starting polymer may have any desirable weight average molecular weight, depending on the final application or use for the hydrogenated product. For example lubricants and additives typically have lower Mw's than do films, fibers, articles and the like. For high strength applications an Mw of at least about 50,000, preferably at least about 100,000, even more preferably, at least about 300,000, even more preferably at least about 400,000 is preferred.

WO 94/21694 PCT/US94/02720

- 8 -

These various starting polymers and block copolymers may be prepared by methods readily known to those of ordinary skill in the art or the block copolymers may be purchased from various suppliers such as Shell Oil Company.

The hydrogenation of these starting block copolymers may be conducted under the usual conditions known to experts in the art utilizing a wide variety of 10 hydrogenation catalysts on an Group 1 or 2 metal salt support. Preferred metal catalysts include those comprising metals selected from groups 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12 of the Periodic Table of Elements. All references to the periodic Table are to the Table as published in Chemical and Engineering News 63(5), 27, 1985. Transition metal catalysts are further preferred with gold, silver, palladium, platinum, rhenium, nickel, rhodium and chromium, being especially preferred. Examples of preferred catalysts include nickel, copper chromate, molybdenum sulfide, palladium, platinum oxide, copper chromium, oxide and the like.

The catalyst is preferably present in the reaction at a ratio of 0.01 to 10.0g of catalyst per gram of polymer, preferably 2.0 to 4.0, even more preferably 2.0 to 5.0. Depending on the choice of catalyst, the catalyst may be reduced prior to the reaction by methods known in the art. The hydrogenation may take place in any reactor which can withstand the temperature and pressure conditions required above.

The catalyst support may be any Group 1 or 2 metal salt, such as lithium, sodium, potassium, barium, calcium, magnesium or cesium salts, preferably barium or calcium salts, more preferably BaSO₄, CaCO₃, BaCO₃. The catalyst is placed on the support by methods known

WO 94/21694 PCT/US94/02720

- 9 -

in the art. The choice of a particular support will be directed by considerations such as the temperature of the reaction, the Group 1 or 2 metal salt selected and the metal catalyst chosen. For Example, when using a calcium carbonate support slightly higher reaction temperatures are desirable to achieve 99.5% or greater unsaturation while slightly lower temperatures may be used for barium sulfate supports and still achieve 99.5% or greater unsaturation. All the possible variations and reaction manipulations are within the skill of one of ordinary skill in the art to determine.

Hydrogenation pressures are preferably in the range from atmospheric to about 3000 psig, preferably between about 100 to about 1000 psig. The hydrogenation temperature can range from 25°C to the degradation of the polymer although the maximum temperatures are preferably restricted to between 100 and 500°C, preferably 100 and 200°C. Reaction times 20 may be as long as necessary to achieve 99.5% or greater hydrogenation. Preferably these times are in the range of 10 minutes to 124 hours or above, even more preferably 2 to 24 hours.

These reactions are preferably conducted in solution phase. Typical solvents are paraffins, isoparaffins, cycloparaffins, tetrahydrofuran and the like and mixtures thereof. Cyclohexanes, hexanes, tetrahydrofuran and mixtures thereof are preferred.

Choice of solvent may be dictated by the starting polymer. For instance, a solvent that allows a greater radius of gyration is preferred over a solvent that allows only a small radius of gyration. For example, polystyrene in cyclohexane with up to about 10 wt% THF, preferably 5 to 10 wt %, has a greater radius of gyration than polystyrene in cyclohexane alone. The

WO 94/21694 PCT/US94/02720

- 10 -

polystyrene in cyclohexane/THF solvent presents the unsaturations so that they are more accessible to the hydrogenation catalysts. Radius of gyration and related phenomena such as solubility parameters and theta solvents, are known parameters and can be measured or calculated by methods known in the art such as light scattering and the like.

The amount of catalyst required for hydrogenation

is a function of the temperature of hydrogenation, of
the degree of hydrogenation desired and within the
skill of one of ordinary skill in the art to determine.
The degree of saturation is proportional to the
reaction temperature, catalyst concentration and mixing

rate. Depending on the degree of saturation required
and the polymer molecular weight all these variables
can be adjusted to modify the reaction rate.

In a typical hydrogenation process an unsaturated 20 polymer, preferably a poly(alkenyl aromatic) or a poly(alkenyl aromatic)/polydiene block copolymer is placed in a solvent, preferably a parraffinic solvent. The solution containing the catalyst is placed in a reactor which is pressured up to 3000 psi, preferably 25 between 100 and 1000 psi, and heated up to 500°C, preferably 100 to 200°C. Depending on the catalyst chosen, the catalyst may be reduced in hydrogen for three hours at 100°C and 100 psig prior to the reaction. The polymer solution is introduced into the 30 reactor and stirred. A preferred catalyst is palladium supported on barium sulfate at a concentration of 2.0 to 5.0 grams, preferably 2.5 grams of catalyst per gram of polymer. The polymer catalyst mixture is then recovered from the reactor and filtered to remove 35 catalyst particles. The hydrogenated polymer is then

. · WO 94/21694 PCT/US94/02720

- 11 -

precipitated in methanol or the like, vacuum dried and stored in air at 0°C.

The degree of hydrogenation is determined from

5 proton NMR (Varian VXR-300) measurements at 25°C using
10% by weight polymer solutions in CDCl₃ with TMS as
the internal reference. Upon hydrogenation the
resonances associated with olefinic and aromatic
protons are converted to aliphatic and cyclic protons.

10 Thus the saturation efficiency can be measured by
analyzing the remaining olefinic and aromatic proton
resonances.

The hydrogenated polymers, preferably the 15 hydrogenated block copolymers produced by this process may then be blended with one or more tackifiers and/or other additives for the formation of adhesives. general, any hydrogenated tackifier will form adhesives with the polymers of this invention. Specific examples 20 of hydrogenated tackifiers include ECR-111 produced by Exxon Chemical Company which is a hydrogenated aliphatic hydrocarbon resin with a Tg of about 35°C, ECR-327 and ECR-143-H which are hydrogenated liquid aliphatic tackifier resins with Tg's of -14°C and -26°C 25 respectively. Tackifiers that are hydrogenated cyclics may also be utilized, however, hydrogenated linear or branched tackifiers are preferred. The tackifiers are preferably present at 50 phr, preferably 100 to 175 phr, even more preferably 100 to 150 phr, even more 30 preferably 110 to 130 phr.

The adhesives can then be coated, laminated or applied onto a substrate, such as wood, paper, polyolefins, film, metal, paper, glass, ceramics and the like.

· . WO 94/21694 PCT/US94/02720

- 12 -

Other oils, plasticizers and processing aids may be added to the tackified composition as well. For example, antioxidants, such as Irganox 1010, fillers, thickeners, flow assist agents, silicon, oils, white mineral oil, low molecular weight polyolefins such as polybutene and chlorinated aromatics such as bisphenyls, and other additives known in the art can be added.

10 EXAMPLES

In the following Examples:

Molecular weight was measured by Gel Permeation 15 Chromotography.

Viscoelastic properties (modulus, tensile strength, elongation at break) were determined using a rheometric solids analyzer (RSA2) operated in the 20 dynamic mode (0.01 < omega < 100 rad/s with a 0.5 mm shear sandwich test fixture). Glass transition temperatures were determined by measuring G' at a fixed frequency. The Tg associated with the glassy component was determined by measuring G' at a fixed frequency 25 (omega_{DS}- $_D$ -i = 0.08 rads/s, omega_{DVCh}- $_{DeD}$ = 1 rad/s) and 2% strain amplitude while slowly heating 1°C per minutes the specimens in nitrogen. The order disorder transition temperature was measured by measuring G' at a fixed frequency and strained amplitude. Frequency 30 equals 0.5 rads/s, 2% strain while slowly heating 1°C/min. Glass transition temperatures for the homopolymers and block copolymers were also determined by differential scanning calimetry (DSC) using a Perk and Elmer system DSC7. The 15 mg test specimens were 35 heated fr m 60 to 200°C at a rate of 10°C per/min and hydrogenation was determined using the method described -WO 94/21694 PCT/US94/02720

- 13 -

above. Gel permeation chromotgraphy (GPC) traces were obtained from a Waters 150c instrument fitted with phenoge columns operated at 25°C with tetrahydrofuran as the mobil phase. Calibration was accomplished using 5 monodisperse standards from Goodyear Chemical Co. (PI, PS-PI) and Pressure Chemical Co. (PS).

Example 1

10 0.47 grams of polyisoprene-polystyrene block copolymer (25% PS) was dissolved in 60 ml of cyclohexane prepared in a purified argon environment and allowed to dissolve for 12 hours. The 2.5 grams of catalyst on Baso4 per gram of polymer was weighed out 15 and placed in the clean hydrogenation reactor with a stir bar. The reactor was evacuated using a two stage vacuum pump, 15 psig of argon was slowly introduced into the reactor and then evacuated again after which the reactor was baked overnight at 115°C. The reactor 20 was then filled with 100 psig hydrogen at a temperature of 100°C, reducing the catalyst for three hours. 100 psig of hydrogen was evacuated and the polymer solution was introduced into the reactor slowly. psig of hydrogen was slowly added to the reactor and 25 was heated to 100°C with stirring at a rate of 2000 rpm. The reaction was terminated after six hours and the reaction was depressurized. The polymer solution was removed and the catalyst particles were allowed to settle out of the polymer solution. The solution was 30 filtered with a 0.45 micrometer filter, precipitated in methanol, vacuum dried and stored in air at 0°C. Tg of the hydrogenated polystyrene polyisoprene block copolymer (PVCH-PEP) was measured to be 141°C. Other data are reported in table 1.

- 14 -

Example 2

The procedure of example 1 was followed, except
that the starting block copolymer was polystyrene 1,4cisbutadiene. Greater than 99% unsaturation was
achieved. Other data are reported in table 1.

Example 3

10

A triblock copolymer of styrene-isoprene-styrene (19 wt.* styrene) was hydrogenated according to the procedure of example 1. Before hydrogenation, the block copolymer had an order disorder transition temperature (ToDT) of 215°C. After hydrogenation the hydrogenated polymer had an order disorder of 245°C. Other data are reported in table 1.

TABLE 1

Example	Wt% PS	Tg (DSC) (*C)	Mw	TODT (°C)	% Sat	Modulus (psi)	Tensile Str (psi)	Elong.	Mw/Mn
1 Before H2	25	101	100k	>300					1.06
1 After H ₂	25	141	100k	>300	>99				
2 Before H2	50		14k	100			1	1	1.06
2 After H ₂	50		14k	235	>99		 		
3 Before H2	19	120	120k	215	1	180	1740	1650	1.06
3 After H ₂	19	130	120k	245	>99	363	1547	800	1

- 15 -

Example 4

Various formulations of hydrogenated block copolymers with tackifiers and non-hydrogenated starting block copolymers with tackifiers were blended and then tested for adhesive properties. The table below describes the components and the adhesive properties.

TABLE 2

10							
	FORMULATION, phr				-		
	Vector 4111 (SIS)	100	100				
	Escorez 1310 (S.P.92C)	100			100		100
15	ECR-111 (S.P.65C)		100	100		120	
	Kraton G 1657			100			
	H ₂ Vector 4111 (Ex 3)				100	100	100
20	Tack Response						
	Quick Stick, lb./in.	3.7	2.1	1.3	0.6	1.0	0.5
	Peel Strength, lb./in.	4.8	3.2	2.7	1.8	1.7	1.7
25	Cohesive Properties						
	Holding Power, Hrs. 1000g. Weight 1" x 1"	100+	100+	100+	100+	25	100+
30	SAFT, 'F 1000g. Weight 1" x 1"	226	234	216	242	230	220

Vector 411 is a triblock (polystyrenepolyisoprene-polystyrene) block copolymer having 19 wt% 35 styren and an Mw of about 120,000.

Escorez 1310 is an aliphatic hydrocarbon rein with a MWD of 1.5 and an Mw of 1250 produced by Exxon Chemical Co. with a Tg of 36 degrees C.

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ECR-111 is a hydrogenated aliphatic hydrocarbon resin with a Tg of about 30 degrees C produced by Exxon Chemical Co.

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Kraton G 1657 is a mixture of triblock (SBS) and diblock (SB) polymers with a saturated diene midblock with an Topy between 200 and 230°C.

Example 5

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The procedure of Example 1 was used except the starting block copolymer was varied. The results and conditions are in Table 3.

20

TABLE 3

POLYM	ER W	T% STYRENE	Mn	Mw/Mn '	Tody (°C)
PS-PI		0.5		18,0001.0	6 124
(PVCH	-PEP)	0.5	18,000	1.06	140
PS-1,	4PBD	0.5	14,000	1.06	100
(PVCH	-PE)	0.5	14,000	1.06	235
PS-1,	2PBD	0.5	50,000	1.06	>300
(PVCH	-PEE)	0.5	50,000	1.06	230

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As is apparent from the foregoing description, the materials prepared and the procedures followed relate to specific embodiments of the broad invention. It is apparent from the foregoing general description and the specific preferred emb diments that while forms of th

• . • WO 94/21694 PCT/US94/02720

- 17 -

invention have been illustrated and described, various modifications can be made without departing from the spirit and scope of this invention. Accordingly, it is not intended that the invention be limited thereby.

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. ' WO 94/21694 PCT/US94/02720

- 18 -

CLAIMS

A method for hydrogenating poly(alkenyl aromatic) polymers and poly(alkenyl aromatic)/polydiene block copolymers to 99.5% saturation or more, said
 method comprising:

contacting under reaction conditions a poly(alkenyl aromatic) polymer or a poly(alkenyl aromatic)/polydiene block copolymer having an Mw of 100,000 or more with a metal catalyst supported on an on a Group 1 or 2 metal salt.

2. The method of claim 1 further comprising recovering a hydrogenated polymer having greater than 99.5% saturation and a Mw/Mn of less than 3.

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The method of claim 1 or 2 wherein the poly(alkenyl aromatic) polymer or the poly(alkenyl aromatic) block is a polymer or copolymer of styrene, para-methyl styrene, methylstyrene, dimethylstyrene,
 para-dimethylstyrene, butyl-styrene, t-butyl-styrene, vinylxylene, ethylvinylxylene, isopropylstyrene, vinylnapthalene, or a mixture thereof and wherein the polydiene is a polymer of one or more C4 to C20 dienes, preferably conjugated or alpha-omega dienes.

25

4. The method of claim 1, 2 or 3 wherein the poly(alkenyl aromatic) polymer is syndiotactic, preferably, syndiotactic poly(alkenyl aromatic) polymer is syndiotactic polystyrene.

30

5. The method of any of the above claims, wherein the catalyst metal is a transition metal, preferably gold, silver, platinum, palladium, rhenium, nickel, rhodium or chromium.

. WO 94/21694 PCT/US94/02720

- 19 -

- 6. The method of any of the above claims wherein the support is a barium or calcium salt, preferably BaSO₄, CaCO₃ or BaCO₃.
- 7. A hydrogenated block copolymer comprising hydrogenated poly(alkyl aromatic)/hydrogenated polydiene blocks and having a Mw/Mn of less than 3 and is greater than 99.5% saturated.
- 10 8. The hydrogenated block copolymer of claim 7, wherein the hydrogenated polydiene is a hydrogenated polymer of conjugated dienes, alpha-omega dienes or a mixture thereof or is a hydrogenated conjugated polyene polymer.

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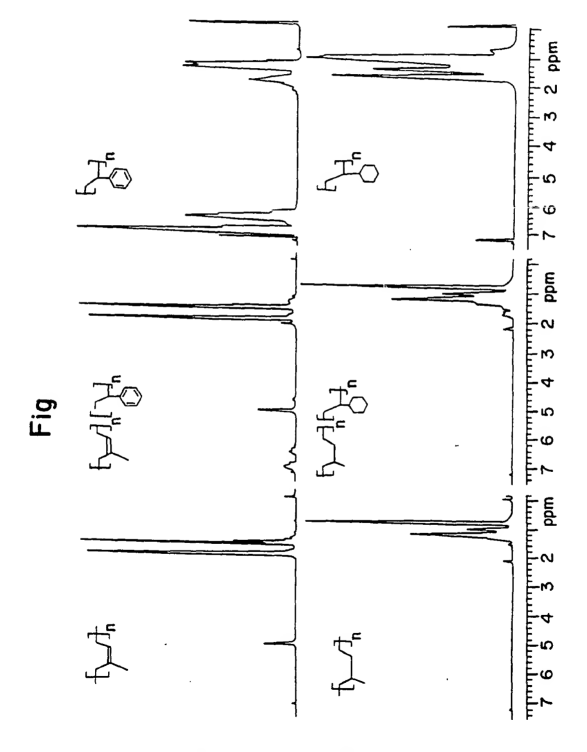
- A 99.5% or greater hydrogenated block copolymer having a Mw/Mn less than 1.5 produced by the process of contacting, under reaction conditions, a polystyrene/polydiene block copolymer with a metal
 catalyst on an Group 1 or 2 metal salt support.
- 10. An adhesive or laminate comprising a 99.5% or greater hydrogenated poly(alkenyl aromatic)/hydrogenated polydiene block copolymer having 25 a Mw/Mn of less than 3 and tackifier.
- 11. The adhesive or laminate of claim 10, wherein the poly(alkenyl aromatic) is polystyrene, the polydiene is polyisoprene or polybutadiene, and the polydiene preferably comprises greater than 40% trans isomers.
 - 12. The adhesive of claim 10 wherein the hydrogenated poly(alkenyl aromatic)/hydrogenated
 p lydiene block copolymer is a hydrogenated (polystyrene/polybutadiene/polystyrene) block copolymer

-WO 94/21694 PCT/US94/02720

- 20 -

having 99.5% or greater saturation and an Mw/Mn less than 1.5 and tackifier.

- 13. A syndiotactic hydrogenated poly(alkenyl5 aromatic) homopolymer, copolymer or block copolymer.
 - 14. A syndiotactic polymer of poly (vinylcyclohexane).



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A. CLASSI IPC 5	FICATION OF SUBJECT MATTER C08F8/04		
According to	o International Patent Classification (IPC) or to both national classific	ition and IPC	
	SEARCHED		
Minimum de IPC 5	ocumentation searched (classification system followed by classification COSF COSC	symbols)	
Documentat	on searched other than minimum documentation to the extent that suc	h documents are inc	luded in the fields searched
Electronic d	ata base consulted during the international search (name of data base i	and, where practical	search terms used)
C. DOCUM	CENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the rele	vant passages	Relevant to claim No.
Y	US,A,3 809 687 (B. ALLISON JR) 7 M see column 1, line 70 - column 2, see column 2, line 49 - line 66; co 1-8	1-14	
Y	EP,A,O 200 029 (BAYER AG) 5 Novembers see page 2, line 21 - page 5, line claims 1-9		1-14
Fu	rther documents are listed in the continuation of box C,	X Patent famil	y members are listed in annex.
"A" docur consist of the consist of the cour which citas of the cour consist of the cour consist of the course of	ment defining the general state of the art which is not idered to be of particular relevance in document but published on or after the international g date ment which may throw doubts on priority claim(s) or his cited to extubits the publication date of another ion or other special reason (as specified) ment referring to an oral disclosure, use, exhibition or a means multiple graph to the international filing date but	or priority date cited to underst invention "X" document of pa cannot be cons involve an invention "Y" document of pa cannot be const document is so ments, such co in the art. "&" document mem	published after the international filing date and not in conflict with the application but and the principle or theory underlying the relevance; the claimed invention dered novel or cannot be considered to naive step when the document is taken alone relevance; the claimed invention idered to involve an inventive step when the minined with one or more other such documentination being obvious to a person skilled ber of the same patent family
1	ne actual completion of the international search	Date of mailing	of the international search report 2 6. 07. 94
	11 July 1994 d mailing address of the ISA European Patent Office, P.B. 5818 Patentlasn 2	Authorized offi	cer
	NL - 2230 HV Ripwith Td. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax. (+31-70) 340-3016	Perme	ntier, W

Information on patent family members

Inter mail Application No PCT/US 94/02720

Patent document cited in search report	Publication date	Patent family member(s)		Publication date	
US-A-3809687	07-05-74	NONE			
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